

Significantly Lower Entropy Estimates for Natural DNA Sequences

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To appear in the Journal of Computational Biology, volume 6, number 1

Abstract

If DNA were a random string over its alphabet $\{A, C, G, T\}$, an optimal code would assign 2 bits to each nucleotide. DNA may be imagined to be a highly ordered, purposeful molecule, and one might therefore reasonably expect statistical models of its string representation to produce much lower entropy estimates. Surprisingly this has not been the case for many natural DNA sequences, including portions of the human genome. We introduce a new statistical model (compression algorithm), the strongest reported to date, for naturally occurring DNA sequences. Conventional techniques code a nucleotide using only slightly fewer bits (1.90) than one obtains by relying only on the frequency statistics of individual nucleotides (1.95). Our method in some cases increases this gap by more than five-fold (1.66) and may lead to better performance in microbiological pattern recognition applications.

One of our main contributions, and the principle source of these improvements, is the formal inclusion of inexact match information in the model. The existence of matches at various distances forms a panel of experts which are then combined into a single prediction. The structure of this combination is novel and its parameters are learned using Expectation Maximization (EM).

Experiments are reported using a wide variety of DNA sequences and compared whenever possible with earlier work. Four reasonable notions for the string distance function used to identify near matches, are implemented and experimentally compared.

We also report lower entropy estimates for coding regions extracted from a large collection of non-redundant human genes. The conventional estimate is 1.92 bits. Our model produces only slightly better results (1.91 bits) when considering nucleotides, but achieves 1.84-1.87 bits when the prediction problem is divided into two stages: i) predict the next amino acid based on inexact polypeptide matches, and ii) predict the particular codon. Our results suggest that matches at the amino acid level play some role, but a small one, in determining the statistical structure of non-redundant coding sequences.

Keywords: *DNA, Entropy, Expectation Maximization (EM), Data Compression, Context Modeling, Time Series Prediction.*

1 Introduction

The DNA molecule encodes information which by convention is represented as a symbolic string over the alphabet $\{A, C, G, T\}$. Assuming each character (nucleotide) is drawn uniformly at random from the alphabet, and that all positions in the string are independent, we know from elementary information theory [Cover and Thomas, 1991, Bell et al., 1990] that an optimal code will devote 2 bits to representing each character. This is the maximum entropy case. DNA may be imagined to be a highly ordered, purposeful molecule, and one might therefore reasonably expect statistical models of its string representation to produce much lower entropy estimates, and confirm our intuition that it is far from random. Unfortunately this has not been the case for many natural DNA sequences, including portions of the human genome.

One example is the human retinoblastoma susceptibility gene containing 180,388 nucleotides, and referred to as HUMRETBAS. The alphabet members do not occur with equal frequency in this string, and accounting for this yields an entropy estimate of roughly 1.95 bits. This is perhaps not surprising since such single character models are the weakest in our information theory arsenal. A logical next step taken by several investigators focuses instead on higher order entropy estimates arising from measurements of the frequencies of longer sequences. For natural languages (*e.g.*, English) this step typically leads to significantly lower entropy estimates. But the best resulting estimate for HUMRETBAS is roughly 1.90 bits [Mantegna et al., 1993], still not impressively different from our 2-bit random starting point. This may be something of a surprise, since such models reflect such known DNA structure as $\%(C + G)$ composition and CG suppression. But these known effects have little impact on the entropy of DNA sequences as a whole.

One may view DNA as a time series, and these higher order models as predictors of the next nucleotide based on its immediate past. With this interpretation the simple 1.95 bit model relies on none of the past. The 1.90 bit result is then even more surprising since it implies that knowledge of the immediate past reduces the entropy estimate by a mere 0.05 bits.¹ Data compression techniques such as Lempel-Ziv (LZ) coding [Ziv and Lempel, 1977] may be viewed as entropy estimators, with LZ corresponding to a model that predicts based on a historical context of variable length. It “compresses” HUMRETBAS to 2.14 bits per character², which is actually worse than the flat random model we started with. The authors’ earlier efforts implemented other advanced modeling techniques but failed to obtain estimates below approximately 1.88 bits.

In this paper we describe a new statistical model, the strongest we are aware of, for naturally occurring DNA sequences. The result is lower entropy estimates for many sequences. Our experiments include a wide variety of DNA, and in almost all cases our model outperforms the best earlier reported results. For HUMRETBAS, the result is approximately 1.7 bits per character — 0.25 bits improved from our 1.95 bit starting level. The standard higher order estimate of 1.90 bits is only 0.05 bits lower, so in this sense our result represents a five-fold improvement. In every case our model outperforms the most common techniques. These results are significant because of our model’s consistent and frequently substantial advantage over earlier methods.

So far we have described the quest for lower entropy estimates as a rather pure *game*, *i.e.*, to predict better an unknown nucleotide based on the rest of the sequence. There are both conceptual and practical reasons to believe that this is an interesting pursuit and that our experimental results are of some importance. Conceptually, the pursuit of better models for data within some domain corresponds rather directly to the discovery of structure in that domain. That is, we model so that we can better understand. Practically, statistical modeling of DNA has proven to be effective in

¹This gap may be thought of as the *mutual information* between the nucleotide being predicted, and its past.

²As implemented in the UNIX (registered trademark of X/Open Company, Ltd.) compress utility.

several problem areas, including the automatic identification of coding regions [Lauc et al., 1992, Cosmi et al., 1990, Cardon and Stormo, 1992, Farach et al., 1995, Krogh et al., 1994], and in classifying unknown sequences into one of a discrete number of classes. The success of such methods ultimately rests on the strength of the models they employ.

In Section 2 we discuss the interpretation of entropy estimates obtained by methods such as ours, and by other approaches such as data compressors. The definition of entropy in the context of DNA nucleotide prediction is made more precise, and we discuss entropy estimates for coding regions.

A detailed exposition of our model is presented in Section 3, but we first introduce and motivate it here in general terms. Conventional fixed-context models focus on a trailing context, *i.e.*, the immediately preceding nucleotides. Longer contexts reach farther into the past, and might reasonably be expected to result in stronger models. However as context length increases, it becomes increasingly unlikely that a given context has *ever* been seen. This problem has led to the development of variable length context language models [Bell et al., 1990], which use long contexts when enough earlier observations exist, and otherwise use shorter ones. Since as earlier noted, the short-term behavior of many natural DNA sequences is so nearly random, these more advanced variable length schemes do not push farther into the past, except in the unusual event of a long exact repeat.

The basic idea behind our model is to push farther into the past by relaxing the exact-match requirement for contexts. Our formalization of this idea in Section 3 is one of the main contributions of this paper.

The metric for determining quality of match is primarily a simple Hamming distance, but in the interests of exploring the use of additional biological knowledge, three other metrics were evaluated in Section 6. We also describe a two-stage model for coding regions in which an amino acid is first predicted based on earlier amino acids, followed by the prediction of a particular codon. Section 7 contains a discussion that suggests why the high entropy levels observed for coding regions may not be so surprising after all, and explores the use of entropy estimation for the classification of sequences.

2 Entropy Estimation and Data Compression

The term *entropy estimate* can be somewhat vague. This section continues our introduction of the term by clarifying its various senses as they relate to DNA, and its relationship to data compression and prediction. We include the section in part to answer some of the most frequently asked basic questions about this work. Our discussion will first focus on the sense of *entropy* that corresponds to a stochastic process. Later, we describe another equivalent sense more suited to sequences of finite extent. The *entropy rate* of a stochastic process $\{X_t\}$ is defined as:

$$\lim_{t \rightarrow \infty} \frac{1}{t} H(X_1, X_2, \dots, X_t) \quad (1)$$

where this limit need not in general exist, and H denotes the information theoretic entropy function (see [Cover and Thomas, 1991]). Given full knowledge of the process, and the limit's existence, the entropy rate is a well-defined attribute of the process. But we know very little of the process underlying the generation of natural DNA, and can merely observe the outcome, *i.e.*, the nucleotide sequence. Given a predictive model \mathcal{X} , and a long DNA sequence $\{x_t\}$, we are therefore content to consider:

$$\lim_{t \rightarrow \infty} \frac{1}{t} \log_2 P(x_t | x_1, \dots, x_{t-1}, \mathcal{X}) \quad (2)$$

This may be thought of as the average bits per symbol required to code increasingly long observation sequences. If our model matches Nature perfectly, Nature’s process is in some sense well-behaved,³ and our sequence is of infinite length, then this limit matches the entropy rate. Of course: i) our model does not match Nature, ii) there is no reason to believe Nature is in the required sense well-behaved, and iii) we can observe only finite DNA sequences. It is in this sense that one is computing an *entropy estimate* when an imperfect model is applied to a finite empirically observed sequence.

Given a data compression program, Equation 2 may therefore be approximated by dividing the number of bits output by the number of symbols (nucleotides) input. We will refer to computations like this as *purely compressive* entropy estimates. Most existing compression algorithms, when applied to natural data, do in fact frequently approach a limit in bits/character — their estimate of the source’s entropy. Unless the sequence being compressed is very long, the purely compressive estimate then overstates the actual entropy. One might then focus only on the code-lengths near the end of the sequence. The risk however is then that the end might just happen to be more predictable and not reflect the overall entropy. For example, the end might, in the case of DNA, contain many long exact repeats of earlier segments. This suggests our second approach, that of *cross-validation* entropy estimates. Here the sequence is first partitioned into N equal size segments. The entropy of each segment is then computed by coding it having first trained the model using the remaining segments. The reported entropy estimate is the average of these N estimates. One can imagine that each of these estimates moves one segment to the end of the sequence, codes it, and averages code-lengths for the final segment only. As N increases, the individual cross-validation estimates may be plotted to visualize the instantaneous entropy rate as a function of position in the sequence. Several of these plots are presented in Section 5.

In what follows, we will switch without harm between a compressive viewpoint, that of stochastic prediction in which the objective is to predict the next nucleotide, and generative stochastic modeling in which one imagines the data to emanate from a particular process.⁴

3 Algorithms

In this section we further motivate our model and then describe it in formal terms.

3.1 Motivation

That natural DNA includes near repeats is well known, and in [Herzel, 1988] the statistics of their occurrence are discussed. We measure *nearness* using ordinary Hamming distance, *i.e.*, the number

³It is sufficient for example that the process be stationary and ergodic — see the Shannon-McMillan-Breiman theorem.

⁴This is possible because a generative process gives rise to a prediction, and a prediction can be converted to a code via arithmetic coding. Starting from a data compressor, we can think of the bits output for a single input symbol, as \log_2 of the probability that a corresponding predictor will predict, or generator will generate that symbol. Unfortunately this isn’t exactly correct for all data compression schemes, *e.g.*, for Lempel-Ziv coding. The heart of the problem is that these codes consider only the greedy parse of the sequence. By this we mean that if one sums over all sequences of length L their inferred probability $2^{-\text{code-length}}$, one will not obtain unity. In most cases the sum is strictly less than unity and the inferred probabilities may be thought of as proportional to probabilities. The result is that entropy estimates are increased by a positive additive constant. In practice however, the discrepancy is modest and one may without harm think of codes as corresponding to probabilities.

of positions at which two equal length strings disagree. Given a target string S , it is then natural to wonder how many h -distant matches exist in a string T . As remarked earlier, DNA seems very nearly random over short distances. Assuming uniform randomness it is easily shown that we expect to see $|T| \cdot 3^h / 4^{|S|} \cdot \binom{|S|}{h}$ h -distant matches. Natural DNA sequences depart markedly from this behavior, providing one part of the motivation for our model. For example (Table 1), using HUMRETBLAS, which consists of 180,388 nucleotides, and a randomly chosen substring of length $w = 20$ from the first $\frac{1}{8}$ of the sequence, one expects 8.61×10^{-6} Hamming distance $h = 1$ matches in the last $\frac{7}{8}$ of the sequence under the random model. We actually see 0.143 matches (see Table 1, column 3), where again, this is an average over all length 20 targets in the last $\frac{7}{8}$ of HUMRETBLAS. When such matches exist, examination of the following nucleotide yields a prediction that is correct 83.77% of the time. That these matches, when present, lead to good predictions is a second motivating factor. Unfortunately such Hamming distance 1 matches occur for only 7.2% of the positions in HUMRETBLAS, so alone, this effect is unlikely to lead to much better predictions on average. But if one again refers to Table 1, roughly $\frac{1}{3}$ of the positions have an earlier match at distance 4, and at even this distance, predictions based on past matches are correct 45.91% of the time. The potential effectiveness of predictions based on matches at all distances is our final motivating factor. The objective, then, is to combine these weak information sources to form a prediction. Also, a version of Table 1 may be made for multiple windows. Our model combines all of this information and makes a single prediction. The use of near matches for prediction is certainly consistent with an evolutionary view in which Nature builds a genome in part by borrowing mutated forms of her earlier work.

One may view each row of Table 1 (really each Hamming distance) as corresponding to a predictive expert, $p_{w,h}(b|S,T)$. The prediction of the Hamming distance 2 expert, $p_{20,2}(b|S,T)$ is formed by examining all matches to the past exactly this distance from our trailing context window and capturing the distribution of following nucleotides by maintaining a simple table of counters. The simplest way to combine these experts is by a fixed set of weights that sum to one.

But suppose that while trying to predict a particular nucleotide b using $w = 20$, our past experience T includes no Hamming distance 0 – 3 matches, *i.e.*, the closest past window is distance $h = 4$. This information is known *before* we see the to-be-predicted nucleotide and may therefore influence the prediction. In particular it makes no sense to give any weight to the opinions of the first three experts — in fact their opinion is not even well-defined in this case. Only the 17 experts corresponding to Hamming distances 4 – 20 are relevant. We’re then interested in properly weighting them conditioned on the assumption that the closest match we’ve seen lies at distance 4. In what follows we will refer to this value as *first Hamming*, f . Finally, since we don’t know *a priori* how window size will influence the prediction, our model is formed at the uppermost level by a mixture of models, each considering a fixed window size from some fixed prior set.

3.2 The CDNA Model

Given a finite alphabet Σ and strings $S, T \in \Sigma^*$ we denote the i th symbol of S by $S[i]$, and the substring consisting of its i th through j th symbols by $S[i, j]$. The concatenation of S and T is denoted $S : T$, and for $b \in \Sigma$ we write $S : b$ to denote the concatenation of S with the length one string consisting of b . Next, the length i suffix of S is denoted $\text{suffix}(S, i)$, and assumes the value of the empty-string if none exists. Finally we denote by $\text{match}(S, T, h)$ the set of all Hamming distance h matches of S in T , e.g. $\text{match}(S, T, 3)$ is the set of all substrings of T that when compared with S exhibit exactly 3 mismatches.

At the center of the CDNA model is an indexed family of probability functions $p_{w,h}$ defined by:

Hamming Distance	Expected # Matches	Observed # Matches	% hit at least once	% correct given 1+ hits
0	1.44×10^{-7}	0.387	4.4%	90.91%
1	8.61×10^{-6}	0.143	7.2	83.77
2	2.45×10^{-4}	0.278	11.0	78.19
3	4.42×10^{-3}	0.505	17.6	66.00
4	0.0563	1.184	33.2	45.91
5	0.541	4.148	78.2	35.86
6	4.01	18.759	98.3	31.90
7	24.3	80.69	100.	29.19
8	119.	304.9	100.	28.35
9	475.	986.0	100.	28.09
10	1566.	2713.	100.	27.86
11	4271.	6286.	100.	27.68
12	9609.	12300.	100.	27.53
13	17740.	20172.	100.	27.33
14	26610.	27387.	100.	27.09
15	31932.	30356.	100.	26.86
16	29936.	26813.	100.	26.57
17	21131.	18200.	100.	26.28
18	10566.	8931.	100.	25.94
19	3337.	2826.	100.	25.69
20	500.	434.5	100.	25.43

Table 1: Predicted vs. observed matches (HUMRETBAS, window size 20). Contexts of length 20 from the first $\frac{1}{8}$ were matched to contexts in the subsequent $\frac{7}{8}$ of the gene.

$$p_{w,h}(b|S, T) = \frac{\mathcal{L} + |\text{match}(\text{suffix}(S, w) : b, T, h)|}{\sum_{\sigma \in \Sigma} \mathcal{L} + |\text{match}(\text{suffix}(S, w) : \sigma, T, h)|} \quad (3)$$

where \mathcal{L} is a Laplace-style *flattening* constant [Laplace, 1825], and is equal to one for the purposes of this paper.

Strings S and T are thought of as time series, and the $p_{w,h}$ provide predictions for next symbol in S and correspond to the *experts* discussed earlier. To limit model complexity we limit the number of experts by restricting w to assume values from some set W .

A simple strategy for combining these experts is to form a weighted sum with the weights totaling unity. But observe that if each term $\text{match}(\text{suffix}(S, w) : \sigma, T, h)$ in Equation 3 is the empty set, then $p_{w,h}$ assigns each $b \in \Sigma$ equal probability $1/|\Sigma|$, and as such is not informative. Following the simple combination strategy then leads to an unnecessarily weak model.

An important contribution of our work is the general idea of ignoring these uninformative experts, and a specific method of doing so. For fixed w we expect more matches as h increases from zero. This is not true in general but is a simplifying assumption we make as modelers. We then focus on the smallest h such that matches exist and ignore all experts before that. Noting that Expert $p_{w,h}$ is uninformative if $\text{match}(\text{suffix}(S, w), T[1, |T| - 1], h)$ is empty, we more formally we define:

$$F(f, w, C, R) = \begin{cases} 1 & \text{if } f = \arg \min_{h \in [1, w]} \text{match}(\text{suffix}(S, w), T[1, |T| - 1], h) \neq \emptyset \\ 0 & \text{otherwise.} \end{cases}$$

The CDNA *prediction for the next character of S* is then the following mixture:

$$P(b|S, T, \Upsilon, \Psi) = \sum_{w \in W} \sum_{f=0}^w \sum_{h=f}^w p_{w,h}(b|S, T) \cdot F(f, w, S, T) \cdot v_{w,f,h} \cdot \psi_w \quad (4)$$

where the $v_{w,f,h}$ and ψ_w are nonnegative weights satisfying $\sum_{w \in W} \psi_w = 1$ and $\sum_{h=f}^w v_{w,f,h} = 1$ for all $w \in W$ and $0 \leq f \leq w$. By Υ and Ψ we denote the parameter values $v_{w,f,h}$ and ψ_w where W is understood to be implicit in them. It follows immediately from the definition of $F(\cdot)$ that Equation 4 is a probability function⁵. The simple combination strategy mentioned earlier involves only two summations, not three. The third, over f , implements our idea of ignoring initial uninformative experts. As a consequence separate mixing parameters are provided for each value of f increasing somewhat the number of parameters in the model. This equation may also be viewed generatively (reading from left to right) as depicted in Figure 1. In this example a window size w is first chosen from the set $W = \{3, 5, 7, \dots, 15\}$. The probability of each choice is shown beside it. Notice that they, and all choices at each node in the diagram, sum to one. In the figure, $w = 9$ is chosen and the next choice is deterministic. Here we've assumed that the closest match to our trailing context of length 9 is distance 3 away. Now given $w = 9$ and $f = 3$ we choose $h \in \{3, \dots, 9\}$ with the probabilities shown. Here we've assumed that $h = 5$ is chosen. Finally, given all the earlier choices, we generate a nucleotide b according to the probabilities shown.

The *online* CDNA predictions of Equation 4 are then combined to form a probability model on strings as follows:

Definition 1 *The CDNA probability of a string S given a reference string T and parameters Υ, Ψ is given by:*

⁵Provided that T is long enough to ensure that some match result is nonempty. Otherwise F will always be zero.

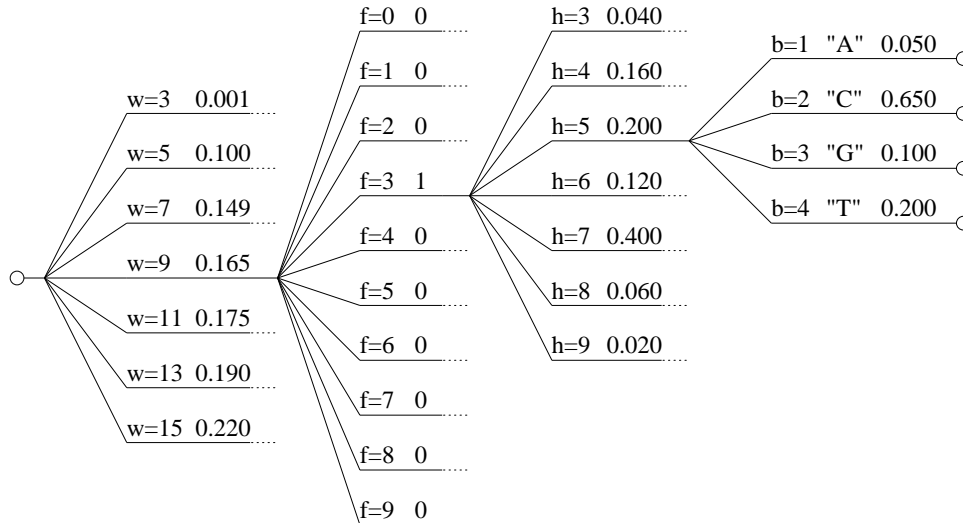


Figure 1: An illustration of the generative branching process that corresponds to the predictive model of this paper.

$$P(S|T, \Upsilon, \Psi) = \prod_{i=1}^{|S|} P(S[i]|S[1, i-1], T, \Upsilon, \Psi)$$

Each prediction of Equation 4 corresponds to a *codelength* $-\log_2 P(b|S, T, \Upsilon, \Psi)$, and with reference to Equation 2, our experimental work that follows averages these codelengths and refers to them as *entropy estimates*.

3.3 The CDNA Learning Algorithm

A CDNA model is parameterized by Υ and Φ . Given a reference string T our learning task is to find parameter values that maximize the probability of S given T . That is, evaluate:

$$\arg \max_{\Upsilon, \Psi} P(S|T, \Upsilon, \Psi) \tag{5}$$

We do not consider the question of optimizing W . But optimizing Ψ in some sense corresponds to learning W since this parameter corresponds to weights on each window size considered. To effect this optimization we apply the Baum-Welch algorithm for Hidden Markov Models [Baum and Eagon, 1967, Baum et al., 1970, Baum, 1972, Poritz, 1988], which may be viewed as an instance of Expectation Maximization (EM) — a later rediscovery [Dempster et al., 1977, Redner and Walker, 1984] of essentially the same algorithm and underlying information theoretic inequality. This approach requires a starting point (in our experiments uniform distributions) and climbs to a local maximum.

We have not considered the complexity of the CDNA global optimization problem but remark that the general problem of globally optimizing directed acyclic graph-based models is known to be NP-complete (see [Yianilos, 1997, p. 32]). Despite our model's simple graphical structure it does exhibit multiple local maxima — but our experience based on trying many random starting points

is that these are very close to one another. Our empirical conclusion is that the uniform starting point is acceptable for the DNA problem domain.

The Baum-Welch/EM parameter reestimates for $v_{w,f,h}$ and ψ_w are given by:

$$v'_{w,f,h} = \frac{Ev_{w,f,h}}{\sum_{i=f}^w Ev_{w,f,i}} \quad , \quad \psi'_w = \frac{E\psi_w}{\sum_{i \in W} E\psi_i}$$

resulting in a new parameter set Υ', Ψ' where

$$Ev_{w,f,h} = \sum_{i=1}^{|S|} \frac{p_{w,h}(S[i]|S[1, i-1], T) \cdot F(f, w, S[1, i-1], T) \cdot v_{w,f,h} \cdot \psi_w}{P(S[i]|S[1, i-1], T, \Upsilon, \Psi)} \quad (6)$$

$$E\psi_w = \sum_{f=0}^w \sum_{h=f}^w Ev_{w,f,h} \quad (7)$$

As a result of Baum-Welch/EM theory we then have

Proposition 1 $P(S|T, \Upsilon', \Psi') > P(S|T, \Upsilon, \Psi)$ unless $\Upsilon' = \Upsilon$ and $\Psi' = \Psi$.

Our implementation differs from the equations above in two respects: 1) it replaces the summation over string positions in Equation 6 with a random sample of positions — with almost no effect on the result, and 2) a small constant (10^{-10}) is added to each $Ev_{h,j,w}$ and $E\psi_w$ term prior to reestimation — to deal with the possibility that some reestimated parameters may not otherwise be well defined. We stress that this is not done for the same reason that one might use Laplace’s rule (add one to event counts) to flatten the prediction of a simple frequency-based model. Such techniques are basically a remedy for data sparseness. We require no such flattening since our model is itself a mixture of many experts, including some for which plenty of data is available. Experiments confirm our intuition that additional Laplace-style flattening is in fact harmful.

4 Experimental Data

The DNA sequences were chosen to pass the following criteria: sufficient length to support this type of entropy estimation method, inclusion of a wide variety of species and sequence types to evaluate the generality of the method, and inclusion of sequences used to benchmark other published methods. All sequences are from GenBank Release No. 92.0 of 18 December 1995. They are: thirteen mammalian genes (GenBank loci HSMHCAPG, HUMGHCSA, HUMHBB, HUMHDABCD, HUMHPRTB, HUMMDBC, HUMNEUROF, HUMRETBLAS, HUMTCRADC, HUMVITDBP, MMBGCXD, MUSTCRA, RATCRYG), a long mammalian intron (HUMDYSTROP), a *C. elegans* cosmid (CEC07A9), seven prokaryote genes (BSGENR, ECO110K, ECOHU47, ECOUW82, ECOUW85, ECOUW87, ECOUW89), a yeast chromosome (SCCHRII), a plant chloroplast genome (CHNTXX), two mitochondrial genomes (MPOMTCG, PANMTPACGA), five eukaryotic viral genomes (ASFV55KB, EBV, HE1CG, HEHCMVCG, VACCG), and three bacteriophage genomes (LAMCG, MLCGA, T7CG).

There has been considerable discussion in the literature of the statistical differences between coding and noncoding regions. To support our study of this issue we assembled several long strands consisting of purely coding or noncoding DNA. These sequences were formed by concatenating coding or non-coding regions from one or more of the above DNA sequences. We consider a region to be coding if either the sense or anti-sense strand is translated; otherwise it is non-coding.

sequence	length	Algorithm						
		UNIX compress	H_1	H_4	H_6	CDNA	bio-com- press-2	CDNA compress
Mammals								
14 Sequences	1029151	2.19	1.98	1.91	1.91	1.67		
Coding only	46457	2.19	1.99	1.94	1.94	1.80		
Noncoding only	982694	2.11	1.98	1.92	1.90	1.74		
HUMDYSTROP	38770	2.23	1.95	1.91	1.95	1.91	1.93	1.93
HUMGHCSA	66495	2.19	2.00	1.92	1.86	0.54	1.31	0.95
HUMHBB	73308	2.20	1.97	1.91	1.92	1.68	1.88	1.77
HUMHDABCD	58864	2.21	2.00	1.92	1.89	1.63	1.88	1.67
HUMHPRTB	56737	2.20	1.97	1.91	1.90	1.69	1.91	1.72
HUMRETBAS	180388	2.14	1.95	1.90	1.90	1.66		1.75
Yeast Chromosome III	315338	2.18	1.96	1.94	1.95	1.90	1.92	1.94
Coding only	211551	2.20	1.97	1.95	1.96	1.92		
Noncoding only	103787	2.19	1.93	1.91	1.92	1.85		
Mitochondria								
MPOMTCG	186609	2.20	1.98	1.96	1.96	1.83	1.94	1.87
PANMTPACGA	100314	2.12	1.88	1.86	1.86	1.81	1.88	1.85
Other Eukaryotes								
CEC07A9	66004	2.20	1.94	1.90	1.92	1.89		
CHNTXX	155844	2.19	1.96	1.93	1.93	1.30	1.62	1.65
Prokaryotes								
7 sequences	784658	2.22	2.00	1.95	1.95	1.92		
Eukaryotic Viruses								
5 sequences	650477	2.15	1.94	1.91	1.91	1.66		
EBV	172281	2.17	1.97	1.93	1.91	1.54		
VACCG	191737	2.14	1.92	1.90	1.90	1.69	1.76	1.81
Bacteriophages								
3 sequences	140739	2.25	1.98	1.94	1.94	1.93		

Table 2: Summary of entropy estimates (bits/nucleotide).

To gather a larger body of coding regions, we obtained a data set of 490 complete human genes. This data set was screened to remove any partial genes, pseudogenes, mutants, copies, or variants of the same gene [Noordewier, 1996]. The resulting sequence contains 484,483 bases and is referred to as our *non-redundant* data set.

5 Experimental Results

Our model’s performance on the sequences described in Section 4 is summarized in Table 2. In some cases our results may be compared directly with estimates from [Grumbach and Tahi, 1994], which are included in the table. Our values for H_4 (the 4-symbol entropy) may be compared with the *redundancy* estimates of [Mantegna et al., 1993] and are in agreement. We have grouped our results by general type (*i.e.*, mammalian, prokaryote, etc.).

The H_1, H_4, H_6 columns contain conventional multigram entropy estimates. The CDNA column reports our model’s cross-validation entropy estimates. Compressive estimates from the BIOPRESS-2 program of [Grumbach and Tahi, 1994] are contained in the following column. Our model’s compressive estimates are given in the table’s final column, CDNA-compress. The compressive estimates are generated by partitioning the sequence into 20 equal segments: s_1, s_2, \dots, s_{20} . The entropy estimate for s_1 , $H(s_1)$ is 2 bits/nucleotide. $H(s_2)$ is calculated using CDNA training on s_1 and testing on s_2 . $H(s_3)$ is calculated from CDNA trained on s_1 and s_2 , tested on s_3 , and so forth. The overall estimate for CDNA-compress is $1/20 \sum_{i=1}^{20} H(s_i)$.

The reported CDNA and CDNA-compress results depend on several model parameters. Their values and a sensitivity analysis are given later in this section. In the mammalian group, “14 Sequences”, and the similarly named lines elsewhere in the table, represent the average of the entropy estimates for all relevant individual sequences, weighted by sequence length.

The conventional multigram entropy results are in general not too informative. It is interesting however to note the variation among sequences in even H_1 , the distribution of individual nucleotides. Only two of these estimates are below 1.9 bits: 1.86 bits for both HUMGHCSA and PANMTPACGA. In the first case CDNA reports 0.54 bits. This is a very repetitive sequence and CDNA is able to exploit this to produce the lowest entropy estimate we observed for any sequence. In this case of PANMTPACGA, nearly all of the redundancy is explained by the unigraph statistics $H_1 = 1.88$. The observed relationship between $\%(C + G)$ and gene density [Dujon et al., 1994] in yeast is reflected in the discrepancy in H_1 between the coding and non-coding region of Yeast chromosome III.

Observe that the H_6 estimate is rarely better than H_4 , and in some cases is markedly worse (a consequence of limited sequence length).

As a point of comparison, entropy rate was also estimated simply by compressing the ASCII representation of the sequence using the UNIX compress utility and dividing the length of the compressed sequence by the length of the uncompressed sequence. Because of its limited dictionary size, the utility cannot be expected to converge to a good entropy estimate for long, complex sequences. For shorter sequences (roughly, sequences not greatly longer than 32,000 nucleotides), this limitation should have no impact. Notice that in all cases, the UNIX compress utility performs worse than no model at all (uniformly random prediction).

To make comparisons among entropy estimation methods clearer, Figure 2 summarizes the results from Table 2 for CDNA, H_6 , BIOPRESS-2, and CDNA-compress. In all cases CDNA outperforms conventional entropy estimates, and in almost all cases by a substantial margin. In many cases the compressive estimates from CDNA (*i.e.*, CDNA-compress) are lower than those of BIOPRESS-2. In several they are comparable, and in only one case (VACCG) does BIOPRESS-2 outperform CDNA-compress by as much as 0.05 bits. It is possible that even this case is due to the offline nature of the current CDNA program which as mentioned above forces us to compute compressive estimates by dividing the sequence into large segments. Finally, we remark that while the table contains results for HUMRETBLAS, this sequence was used in our parameter sensitivity study below.

The CDNA program’s behavior is parameterized by several values:

- The number of EM iterations to perform when learning the model’s parameters based on the training set.
- The set of allowed window sizes (trailing contexts), W .
- The size of the training sample.
- The size of the test sample.

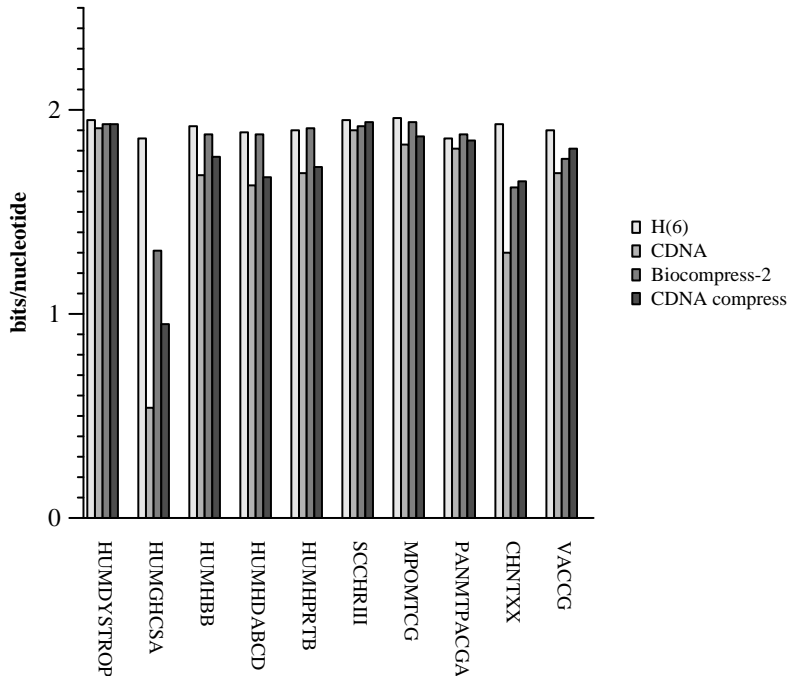


Figure 2: Comparison of results.

- The number of cross-validation segments.
- The random seed that controls all sampling.
- The distance measure to use.

The distance measure is discussed later in Section 6 and for all other experiments is fixed as described earlier. We used 3-way cross-validation to assess parameter sensitivity (see discussion below), and 8-way cross-validation for all experiments. Our test samples were of size 1000 per cross-validation segment for a total of 8000 positions. To assess the variation due to all sampling, we performed 11 different cross-validation estimates using different random seeds. The mean entropy estimate for HUMRETBLAS was 1.6939 bits with a standard deviation of 0.014 bits. Larger samples would result in smaller variance at the expense of computer time.

Figure 3 shows the effect of EM iterations on the entropy estimate for both training and test sets. This figure and the two that follow employ HUMRETBLAS. Based on this analysis, we chose 200 EM iterations for our experiments. Notice that there is little overtraining. Both train and test performance are essentially constant after 150 iterations, and their gap is small.

Figure 4 shows the effect of a single window size on performance. Beyond 25, overtraining is evident. This is to be expected since the number of parameters grows quadratically with window size. For our experiments we selected a mixture of windows of size 2, 4, 6, 8, \dots , 24. Odd values were excluded only to reduce memory requirements for the model. Additional experiments indicated that while mixtures of different window sizes did in general help, the benefit was somewhat small. We did not study this for all sequences and therefore employed a mixture to ensure that the model is as general as possible (in theory the mixture can only help).

Finally, Figure 5 shows the sensitivity of performance on training set size. We used a training set of size 5000 for our experiments because beyond that point, little improvement was evident: train and test performance had essentially converged.

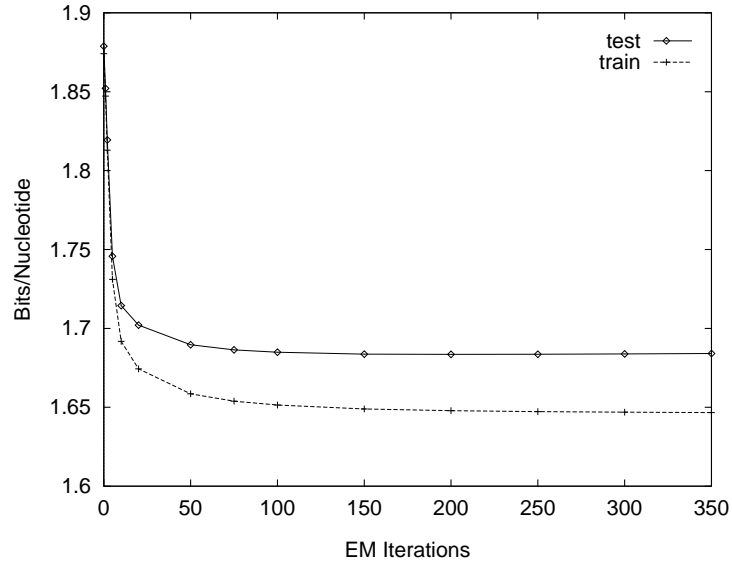


Figure 3: The effect of varying the number of EM learning iterations on performance for HUMRET-BLAS.

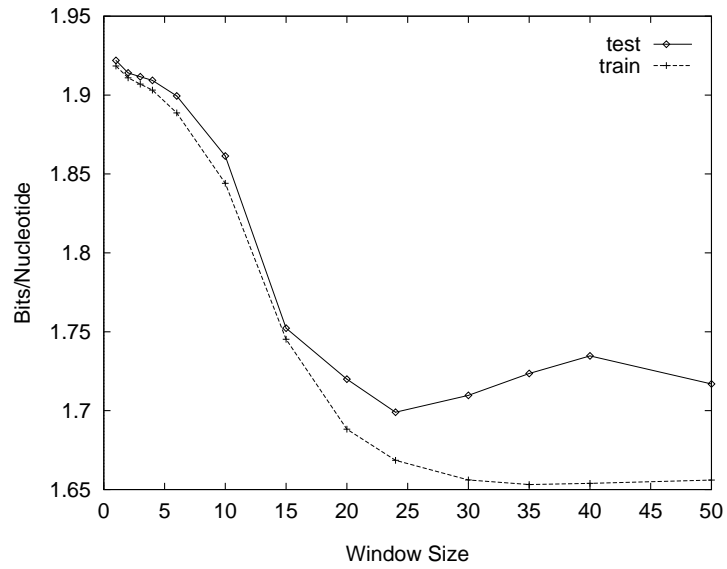


Figure 4: The effect of varying the trailing context size (window) on performance for HUMRETB LAS.

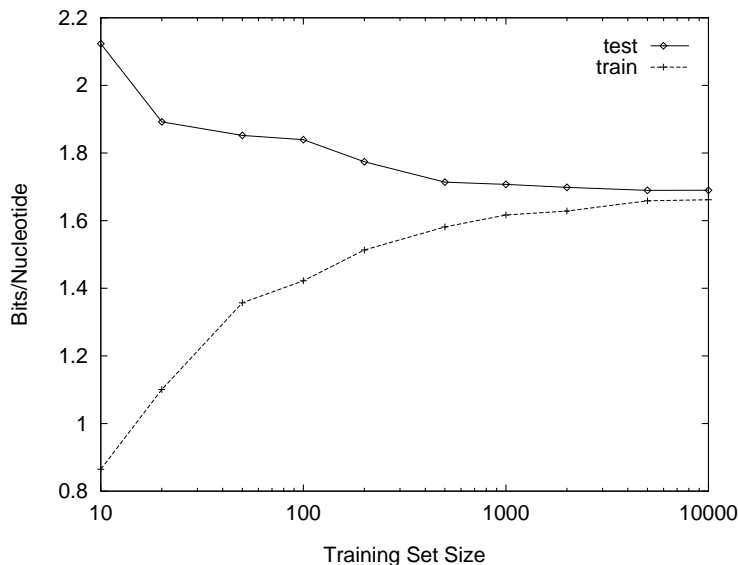


Figure 5: The effect of varying the size of the training set on performance for HUMRETLAS.

Training Set	Test Data	
	Coding	Noncoding
Coding	1.829	2.010
Noncoding	1.949	1.727

Table 3: Mammalian corpus cross-entropies (bits/nucleotide).

Our results so far have consisted of entropy estimates for sequences of a single type. We now present the results of several cross-entropy experiments. By this we mean that the type of the training set is different from that of the test set. In Table 3 mammalian coding regions and non-coding regions form the two types. A substantial cross-entropic gap is apparent. That is, a model trained on the coding sequence might be used to distinguish coding from non-coding based on entropy. The same is true when one trains on the non-coding sequence.

The same experiment produces very different results for yeast as shown in Table 4. While a large gap exists when one trains using the non-coding regions, a *reverse* gap exists in the other case. That is, training on the coding regions assigns shorter codes to the non-coding regions, than to itself. While this effect is weak, it suggests that sequences from the coding region may be *echoed* in the non-coding region.

Cross-species entropy estimates are interesting in general, but our one example shown in Table 5 shows little interesting structure. That is, yeast coding regions teach nothing about human coding

Training Set	Test Data	
	Coding	Noncoding
Coding	1.941	1.918
Noncoding	1.982	1.831

Table 4: Yeast chromosome III cross-entropies (bits/nucleotide).

Training Set	Test Set	Entropy
Mammalian Coding	Yeast Coding	2.026
Mammalian Noncoding	Yeast Noncoding	1.927
Yeast Coding	Mammalian Coding	2.033
Yeast Noncoding	Mammalian Noncoding	1.955

Table 5: Yeast/Mammalian cross-entropies (bits/nucleotide).

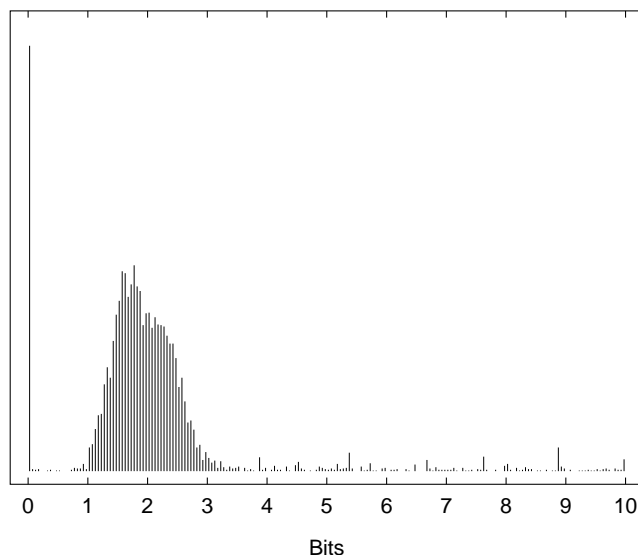


Figure 6: Histogram of nucleotide code-lengths for EBV.

regions and visa versa. In fact, the entropy estimates are in both cases worse than random. In the non-coding case, a weak relationship exists, but is easily explained by agreement on the low order multigraphic statistics of the sequences, *e.g.*, H_4 .

The entropy estimates we have discussed reduce to a single number the model's performance on a sequence. It represents the average code-length but says nothing about their distribution. The histograms of figures 6,7, and 8, depict this distribution for the EBV virus, the human gene HUMRETBLAS, and the yeast chromosome SCCHRIII. They present an interesting visual *signature* of a sequence. Trimodality is apparent in all three, but is present to a varying extent.

The first mode, most prevalent in our viral example, corresponds to near-zero code-lengths that correspond to long exact or near-exact repeats. In the viral case, this peak is particularly narrow because while the sequence is known to contain several long exact repeats, it lacks the full range of near repeats of various lengths found in sequences such as HUMRETBLAS. The middle mode corresponds to nucleotides having a long context that matches well with some other segments from the sequence. In this case short codes are assigned so long as these matches predict well the following nucleotide. The rightmost mode, particularly pronounced in SCCHRIII and less so in HUMRETBLAS, correspond to codes well over 2 bits. Here the model had a strong prediction, but was wrong.

The distribution of code-lengths displayed in these histograms says nothing about how code-lengths are distributed with respect to position in the DNA sequence. To visualize this we have plotted in Figures 9 and 10 the code-lengths at 8000 random positions for HUMRETBLAS and SCCHRIII. It is apparent that the distribution is not homogeneous and presents another interesting

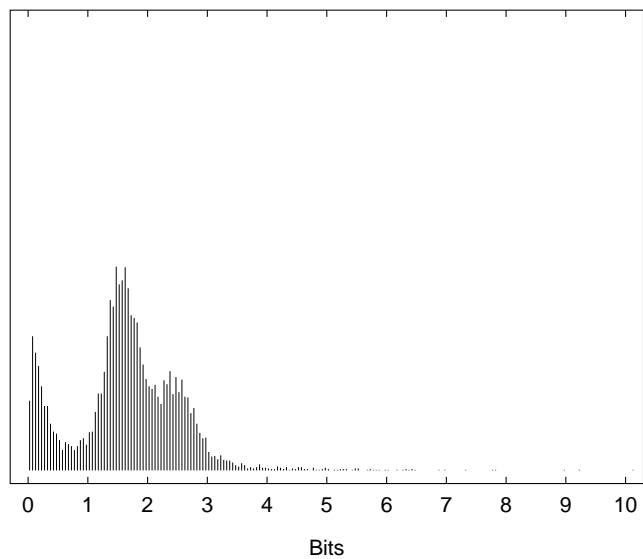


Figure 7: Histogram of nucleotide code-lengths for HUMRETLAS.

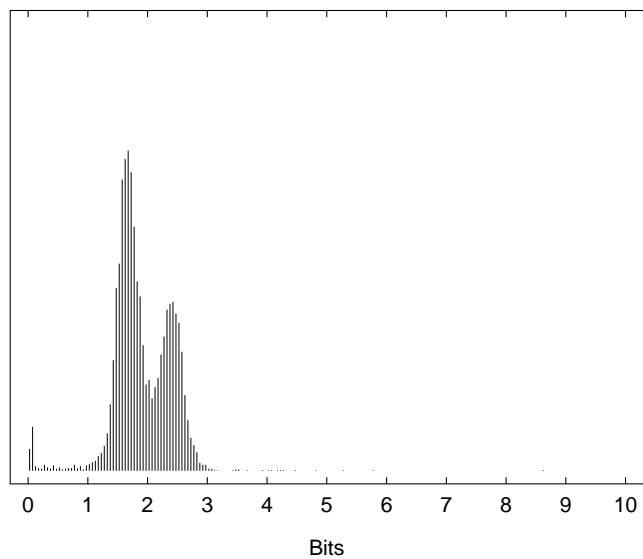


Figure 8: Histogram of nucleotide code-lengths for SCCHRIII (yeast).

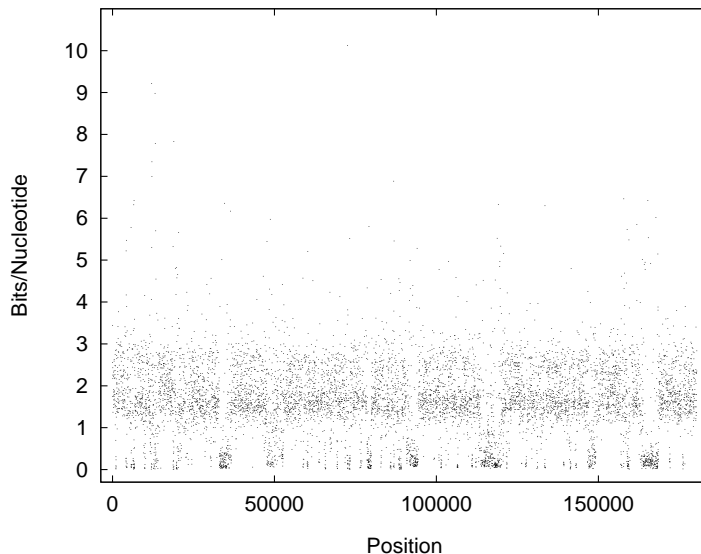


Figure 9: The variation of code-length with position for HUMRETLAS.

visual signature of the sequence. The distribution for HUMRETLAS is shown at a closer scale (Figure 11) where finer structure is revealed. The modality noted in the histograms is also apparent in these plots.

6 Alternative Metrics

Our model exploits near matches to form a prediction, and its performance therefore depends on precisely what is meant by *near*, *i.e.*, the distance metric used to judge string similarity. The simplest notion of string distance relevant to our domain is simple Hamming distance: the number of mismatching nucleotides in two strings of equal length. We will refer to this as the *nucleotide-sense* metric.

Domain knowledge suggests that we also compare the strings after reversing one, and complementing its bases, so that matches may occur between the sense and anti-sense strands of DNA. Both distances are computed and the smaller one used. This is our *nucleotide-both* metric, and was used in all experiments from earlier sections.

A common misconception is that coding regions should compress well because three nucleotides together (64 possibilities) code for only one of 20 amino acids. This reasoning is flawed because 61 of the 64 possibilities represent valid codes for amino acids: the genetic code is degenerate. Thus the maximum entropy level per nucleotide for coding regions is not $\log_2 20/3 \approx 1.44$ bits, but rather is $\log_2 61/3 \approx 1.977$ bits. In fact, it has been observed by several authors that coding regions are less compressible than non-coding regions (*e.g.*, [Salamon and Konopka, 1992, Farach et al., 1995]). So it is clear that two sequences that code for the same polypeptide may nevertheless have large Hamming distance.

Our first approach to making use of the degeneracy of the genetic code is to replace the Hamming distance in the match function described in Section 3.2 with an *amino-Hamming distance*. This distance is defined to be the smallest Hamming distance, examined over all three possible reading frames, between the amino-acid translations of the two substrings. Using this modified distance either without or with matches to the antisense strand yields the *amino-sense* and *amino-both*

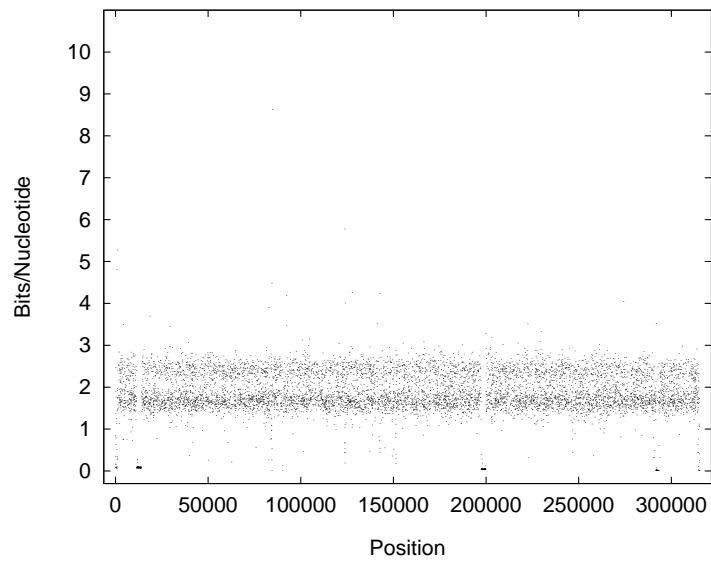


Figure 10: The variation of code-length with position for SCCHR111 (yeast).

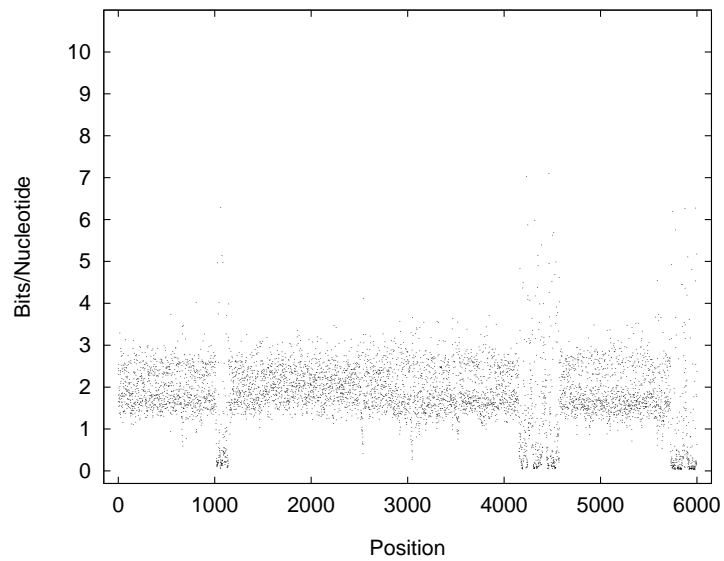


Figure 11: The variation of code-length with position for HUMRETBLAS in the first 6,000 positions.

sequence	nucleotide		amino acid	
	sense	both	sense	both
Mammals				
Coding only	1.830	1.830	1.882	1.895
Noncoding only	1.750	1.727	1.795	1.769
HUMGHCSA	0.525	0.510	0.550	0.547
HUMRETBLAS	1.748	1.686	1.799	1.731
SCCHRIII	1.904	1.879	1.930	1.926
Coding only	1.947	1.941	1.966	1.970
Noncoding only	1.879	1.831	1.889	1.850
Prokaryotes				
7 sequences	1.944	1.939	1.977	1.977

Table 6: Effect of metric on entropy estimate (bits/nucleotide).

metrics respectively.

Selected experiments comparing these four metrics are shown in Table 6.⁶ Entire sequences as well as their noncoding and coding regions were considered. The *nucleotide-both* metric performs best, and was therefore used for the experiments reported in earlier sections.

The amino metrics never outperformed their nucleotide counterparts. This is not so surprising for entire sequences, or noncoding regions, but even for coding regions, the amino metrics perform somewhat worse. Their poor performance is explained by the fact that they distinguish fewer discrete distances, and are always blind to the exact nucleotide structure of the sequence, even in the positions immediately before the base to be predicted.

Our second approach separates the prediction problem into two stages. An amino acid is first predicted using CDNA based on inexact matches, where the sequence is viewed at the amino acid level, *i.e.*, has alphabet size 20. The model’s second stage predicts a particular codon, conditioned on the already predicted amino acid, and on earlier positions which are now viewed at the nucleotide level. The codon entropy estimate is then the sum of the entropies of these two stages: $H(C) = H(A) + H(C|A)$. Dividing $H(C)$ by three then gives a per-nucleotide entropy which may be compared with earlier results.

Table 7 gives the results of several models on our *non-redundant* data set, including multigram statistics and CDNA applied to both nucleotide and amino-acid representations. In the case of CDNA, 2000 training samples were used. The last line of the table employs what amounts to *maximal* cross validation, where a sequence element is predicted using everything else in the sequence.

It is apparent that our non-redundant collection of coding material is harder to model than the entire genes considered in earlier sections. Nevertheless we are able to improve the 1.92 multigraphic estimate somewhat to 1.84. It is important to note our non-redundant data set is not a random sample, but rather is closer to the worst case.

7 Discussion

We begin by discussing the question “what entropy level do we expect and why?” As remarked earlier the apparent incompressibility of DNA is surprising, but this implies that we expect much

⁶These data were collected using smaller numbers of training and test samples than those presented in Table 2.

Model Description	bits/nuc.
Maximum Entropy	1.98
Nucleotide 6-grams (H_6)	1.92
CDNA nucleotide	1.91
Amino 2-grams ($H_2(A)$) + Codon-predict ($H(C A)$)	1.89
CDNA amino (8-way cross-validation) + Codon-predict	1.87
CDNA amino (max. way) + Codon-predict	1.84

Table 7: Summary of modeling results (bits/nucleotide) for coding regions extracted from a *non-redundant* collection of 490 human genes .

lower entropy estimates. We submit that the essential cause of our surprise is that simple statistical models, which perform so well in other cases, fail to discover significant structure in DNA. Our work demonstrates that DNA is more predictable than had previously been established, but falls far short of the compression levels easily achieved for natural language. This suggests that local structure parallel to “words” and “phrases” is largely missing in DNA, even when the exact match requirement is relaxed.

We believe that the question ultimately comes down to one of quantifying the magnitude of the functional degeneracy of polypeptides. That is, how many different sequences are essentially equivalent from the standpoint of natural selection? If this degeneracy turns out to be large, then DNA is to its function, as an acoustic waveform is to human speech: small local distortions do little to alter the overall message.

As another illustration of this *counting* view of entropy, suppose for that the total length of every human’s DNA is an identical value N . Now let M be the number of nucleotide sequences of this length that code for viable humans. Then each human genome may be coded using $\log_2 M$ bits and the per-character entropy is then $(\log_2 M)/N$. This is, on the one hand, an elegant and satisfying definition, but on the other, it is difficult to imagine computing it. We can, however, imagine injecting domain knowledge at a less grandiose scale. For example, models of a coding region might condition their predictions on knowledge of which polypeptides correspond to plausibly useful proteins. Even inexact knowledge of this relationship could result in lower entropies.

The CDNA model uses only one kind of domain knowledge: that natural DNA sequences include more near repetitions than one expects at random. It is possible that the addition of more domain knowledge will produce lower entropy estimates for coding regions. However, we believe that the actual entropy may be not so far below our current estimates. This is consistent with the view that the coding regions of DNA commonly amount to a highly random string, with only isolated critical regions. If much of large proteins amounts to *scaffolding*, and the bulk of proteins in a genome are large, then we would expect large entropies. By scaffolding we mean weakly constrained structure necessary only to ensure that the protein folds correctly, and that particular active regions are positioned properly in the resulting molecule. There are certainly many genes that code for somewhat small proteins, but since entropy is a per-base expected value, the genes that code for large proteins have proportionally greater affect on our estimate. In our non-redundant collection of human genes, fewer than one third of bases are contained in coding regions of length 1000 or less.

We quantify this conjecture with a crude analysis. If only the hydrophobic/hydrophilic nature of an amino acid were critical, then at most one bit would be required to make an acceptable selection. This is then $\frac{1}{3}$ bit per base. If, except for this structure, the string is random, then the

lowest entropy we would expect is: $1.644 = 1.977 - 0.333$ bits per base. Biological experiments to quantify the sensitivity of proteins of various length to mutation, would then shed light on the actual entropy of coding regions.

We conclude our discussion by remarking on the application CDNA to classification problems. A data compressor may be used as a classifier, and the result interpreted in probabilistic terms. This idea was introduced in the context of bioinformatics by [Loewenstern et al., 1995] where compressive classification was applied to several problems including promoter recognition. To illustrate, suppose one has two long sequences x, y of DNA, the first of some type A and the other of type B . The task is to classify a third strand z . Let $L(s)$ denote the bits output by the compressor when fed sequence s . Then $L(x : z) - L(x)$ may by our remarks above be interpreted as $-\log_2 P(z|x)$. Similarly $L(y : z) - L(y)$ corresponds to $-\log_2 P(z|y)$. At this point one might simply compare to affect classification, or exponentiate yielding probabilities. These can then be combined along with prior class probabilities, resulting in Bayesian classifier built from data compressors. Later work [Loewenstern et al., 1997] applied this general idea to classify short DNA sequences by their x-ray crystallographic structure.

8 Conclusion and Future Work

We have shown that the near repeats in natural DNA sequences may be incorporated into a statistical model resulting in significantly lower entropy estimates. For some sequences our model is the first to detect substantial deviations from random behavior and illustrates the importance of inexact string matching techniques in this field. It is entirely possible that very different results will be obtained, particularly for coding regions, when much more DNA is available for analysis.

It should be noted that H_1 entropy estimates include the known effect of $\%(C+G)$ [Gatlin, 1972] on entropy estimation, and BIOCOMPRESS-2 includes the also known effect of long exact repeats and exact complement repeats [Herzel et al., 1994]. CDNA's generally superior performance indicates that DNA possesses more structure which may be exploited.

Several notions of distance were evaluated and the best performance resulted from considering both Hamming distance to reversed and complemented targets, as well as standard Hamming distance, then selecting the minimum. We also described two-stage models crafted especially for coding regions, which perform better than our basic model on coding sequences.

Future work will consider stronger models exploiting additional effects. The current assumption that each prediction event is independent of the last may be dropped and a temporal dependence component added to the model. Next the metric may be enhanced to notice not only the number of mismatches, but also where they occur. Other structural changes are possible and the inclusion of additional domain knowledge in some form is a particularly interesting direction. Also, there are entirely different approaches one might take to the problem of combining experts.

We will also investigate fully *on-line* models that learn as each symbol is processed and use a more sophisticated data structure to greatly reduce the time needed to compute match sets.

Acknowledgments

We thank Martin Farach, Haym Hirsh, Harold Stone, and Sarah Zelikovitz for helpful comments on earlier drafts of this paper.

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